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PTO/SB/08a (07-09)

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<p>Substitute for form 1449A/PTO</p> <h2 style="text-align: center;">INFORMATION DISCLOSURE STATEMENT BY APPLICANT</h2> <p><i>(Use as many sheets as necessary)</i></p>				Complete if Known	
				Application Number	10/567,424
				Filing Date	12/9/2008
				First Named Inventor	Mulley et al.
				Art Unit	1649
				Examiner Name	Kolker, Daniel E.
Sheet	1	of	16	Attorney Docket Number	

U.S. PATENT DOCUMENTS

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				First Named Inventor	Mulley et al.
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				Examiner Name	Kolker, Daniel E.
Sheet	2	of	16	Attorney Docket Number	
1386/23					

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	33	Abstracts of Decisions. Decision of a Delegate of the Commissioner of Patents corresponding to an Australian Patent Application No. 18465/01 issued January 29, 2007.			
	34	Alekov et al., "A sodium channel mutation causing epilepsy in man exhibits subtle defects in fast inactivation and activation <i>in vitro</i> ," Journal of Physiology, Vol. 529, No. 3, pgs. 533-539 (2000).			
	35	Andermann, "Multifactorial Inheritance of Generalized and Focal Epilepsy," Genetic Basis of the Epilepsies, pgs. 355-374 (1982).			
	36	Annegers, "The Epidemiology of Epilepsy," The Treatment of Epilepsy: Principles and Practice, Chpt. 11, pgs. 165-172 (1996).			
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	40	Berkovic et al., "Concepts of absence epilepsies: Discrete syndromes or biological continuum?" Neurology, Vol. 37, No. 6, pgs. 993-1000 (June 1987).			
	41	Berkovic et al., "Familial Epilepsies in Israel: Clinical Syndromes and Modes of Inheritance," Neurology, Vol. 54, Suppl. 3, A356, No. P05.063 (April 2000).			
	42	Berkovic et al., "The epilepsies: specific syndromes or a neurobiological continuum?" Epileptic Seizures and Syndromes, Chpt. 5, pgs. 25-37 (1994).			

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	43	Bertrand et al., "Properties of neuronal nicotinic acetylcholine receptor mutants from humans suffering from autosomal dominant nocturnal frontal lobe epilepsy," British J. of Pharmacology, Vol. 124, pgs. 1-10 (1998).		
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	53	Communication pursuant to Rule 46(1) EPC corresponding to European Application No. 04718885.9-2402 PCT/AU2004000295 dated July 14, 2006.				
	54	DATABASE UniProt, "Sodium channel protein type I alpha subunit," XP002313393, retrieved from EBI accession no., UNIPROT: CIN1_HUMAN, Database accession no. P35498. (ABSTRACT)				
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	56	Doose and Baier, "Genetic Aspects of Childhood Epilepsy," Cleveland Clinic Journal of Medicine, Vol. 56, Suppl. Part 1, S101-S110 (1989).				
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	58	Dworakowska and Dolowy, "Ion channels-related diseases," ACTA Biochimica Polonica, Vol. 47, No. 3, pgs. 685-703 (2000).				
	59	Escayg et al., "Mutations of SCN1A, encoding a neuronal sodium channel, in two families with GEFS+2," Nature Genetics, Vol. 24, pgs. 343-345 (April 2000).				
	60	European Patent Office Search Report corresponding to European Patent Application No. 07075566.5 - 2401 dated October 4, 2007.				
	61	Examiner's First Report for Australian Patent Application No. 2004200978 dated May 6, 2004.				
	62	Finkelstein et al., "Use of denaturing gradient gel electrophoresis for detection of mutation and prospective diagnosis in late onset ornithine transcarbamylase deficiency," Genomics, Vol. 7, pgs. 167-172 (1990).				

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	63	Fong et al., "Childhood Absence Epilepsy with Tonic-Clonic Seizures and Electroencephalogram 3–4-Hz Spike and Multispike-Slow Wave Complexes: Linkage to Chromosome 8q24," Am. J. Hum. Genet., Vol 63, pgs. 1117-1129 (1998).				
	64	Fujiwara et al., "Long-Term Course of Childhood Epilepsy with Intractable Grand Mal Seizures," Jpn. J. Psychiatry Neurol., Vol. 46, pgs. 297-302 (1992).				
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	67	Genbank accession number AB093548.	10/16/2002			
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	73	Gene Card for SCNA1 available via uri: <genecards.org/cgi-bin/cardisp.pl?gene=SCN1A> 11/16/2006				
	74	GeneCards output for protein-coding SCN1A, available online from www.genecards.org, pgs. 1-20. 7/23/07				
	75	Gennaro et al., "Familial severe myoclonic epilepsy of infancy: truncation of Na _v 1.1 and genetic heterogeneity," Epileptic Disord., Vol. 5, pgs. 21-25 (2003).				
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	83	Hauser et al., "Incidence of Epilepsy and Unprovoked Seizures in Rochester, Minnesota: 1935-1984," <i>Epilepsia</i> , Vol. 34, No. 3, pgs. 453-468 (1993).		
	84	Hille, "Ionic Channels of Excitable Membranes," 2 nd Edition, pgs. 423 and 434-444 (1992).		
	85	Hirschhorn et al., "A comprehensive review of genetic association studies," <i>Genetics in Medicine</i> , Vol. 4, No. 2, pgs. 45-61 (2002).		
	86	International Search Report for International Application No. PCT/AU2004/000295 dated May 14, 2004.		
	87	Interview Summary corresponding to U.S. Patent Application Serial No. 11/262,647 dated December 5, 2008.		
	88	Interview Summary corresponding to U.S. Patent Application Serial No. 10/806,899 dated November 23, 2009.		
	89	Interview Summary corresponding to U.S. Patent Application Serial No. 10/806,899 dated October 8, 2009.		
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	91	Janz et al., "Do idiopathic generalized epilepsies share a common susceptibility gene?" <i>Neurology</i> , Vol. 42, Suppl 5, pgs. 48-55 (April 1992).		
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	93	Kimura K., "A missense mutation in SCN1A in brothers with severe myoclonic epilepsy in infancy (SMEI) inherited from a father with febrile seizures," Brain Dev., Vol. 27, No. 6, pgs. 424-430 (September 2005).				
	94	Kinzler et al., "Identification of a gene located at chromosome 5q21 that is mutated in colorectal cancers," Science, Vol. 251, pgs. 1366-1370 (1991).				
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	98	Lernmark and Ott, "Sometimes it's hot, sometimes it's not," Nature Genetics, Vol. 19, pgs. 213-214 (July 1998).				
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				Application Number	10/567,424
				Filing Date	12/9/2008
				First Named Inventor	Mulley et al.
				Art Unit	1649
				Examiner Name	Kolker, Daniel E.
Sheet	10	of	16	Attorney Docket Number	
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	107	Moran et al., "Skeletal Muscle Sodium Channel Is Affected by an Epileptogenic β1 Subunit Mutation," <i>Biochem. Biophys. Res. Comm.</i> , Vol. 282, pgs. 55-59 (2001).			
	108	Moulard et al., "Identification of a New Locus for Generalized Epilepsy with Febrile Seizures Plus (GEFS+) on Chromosome 2q24-q33," <i>Am. J. Hum. Genet.</i> , Vol. 65, pgs. 1396-1400 (1999).			
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	110	Mulley et al., "Channelopathies as a Genetic Cause of Epilepsy," <i>Current Opinion in Neurology</i> , Vol. 16, pgs. 171-176 (2003).			
	111	Notice of Allowance corresponding to U.S. Patent Application Serial No. 10/451,126 (Patent No. 7,078,515) dated August 30, 2005.			
	112	Notice of Allowance corresponding to U.S. Patent Application Serial No. 11/263,326 (Patent No. 7,282,336) dated June 18, 2007.			

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	114	Notice of Allowance corresponding to U.S. Patent Application Serial No. 10/806,899 dated January 4, 2010.				
	115	Notification Concerning Transmittal of Copy of International Preliminary Report on Patentability for International Application No. PCT/AU2006/000841 dated January 3, 2008.				
	116	Official Communication of U.S. Patent and Trademark Office corresponding to U.S. Patent Application Serial No. 10/482,834 dated August 7, 2009.				
	117	Official Communication of U.S. Patent and Trademark Office corresponding to U.S. Patent Application Serial No. 10/482,834 dated December 30, 2008.				
	118	Official Communication of U.S. Patent and Trademark Office corresponding to U.S. Patent Application Serial No. 10/482,834 dated April 4, 2008.				
	119	Official Communication of U.S. Patent and Trademark Office corresponding to U.S. Patent Application Serial No. 10/482,834 dated August 2, 2007.				
	120	Official Communication of U.S. Patent and Trademark Office corresponding to U.S. Patent Application Serial No. 10/806,899 dated October 28, 2009.				
	121	Official Communication of U.S. Patent and Trademark Office corresponding to U.S. Patent Application Serial No. 10/806,899 dated May 13, 2009.				
	122	Official Communication of U.S. Patent and Trademark Office corresponding to U.S. Patent Application Serial No. 10/806,899 dated August 19, 2008.				

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				Examiner Name	Kolker, Daniel E.
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	124	Official Communication of U.S. Patent and Trademark Office corresponding to U.S. Patent Application Serial No. 10/806,899 dated November 29, 2006.		
	125	Official Communication of U.S. Patent and Trademark Office corresponding to U.S. Patent Application Serial No. 11/263,326 (Patent No. 7,282,336) dated October 6, 2006.		
	126	Official Communication of U.S. Patent and Trademark Office corresponding to U.S. Patent Application Serial No. 11/262,647 dated April 22, 2009.		
	127	Official Communication of U.S. Patent and Trademark Office corresponding to U.S. Patent Application Serial No. 11/262,647 dated December 5, 2008.		
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				Filing Date	12/9/2008
				First Named Inventor	Mulley et al.
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	133	Ohtsuka et al., "Refractory Childhood Epilepsy and Factors Related to Refractoriness," <i>Epilepsia</i> , Vol. 41, Suppl. 9, pgs. 14-17 (2000).				
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Examiner Signature	/Daniel Kolker/	Date Considered	02/09/2011
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